Food and Drug Administration, HHS

premarket notification procedures in subpart E of part 807 of this chapter subject to the limitations in §868.9.

[47 FR 31142, July 16, 1982, as amended at 61 FR 1120, Jan. 16, 1996; 66 FR 38795, July 25, 2001]

\$868.5580 Oxygen mask.

- (a) *Identification*. An oxygen mask is a device placed over a patient's nose, mouth, or tracheostomy to administer oxygen or aerosols.
- (b) Classification. Class I (general controls). The device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter subject to the limitations in §868.9.

[47 FR 31142, July 16, 1982, as amended at 61 FR 1120, Jan. 16, 1996; 66 FR 38795, July 25, 2001]

§868.5590 Scavenging mask.

- (a) *Identification*. A scavenging mask is a device positioned over a patient's nose to deliver anesthetic or analgesic gases to the upper airway and to remove excess and exhaled gas. It is usually used during dentistry.
- (b) Classification. Class I (general controls). The device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter subject to the limitations in §868.9.
- [47 FR 31142, July 16, 1982, as amended at 61 FR 1120, Jan. 16, 1996; 66 FR 38795, July 25, 2001]

§868.5600 Venturi mask.

- (a) *Identification*. A venturi mask is a device containing an air-oxygen mixing mechanism that dilutes 100 percent oxygen to a predetermined concentration and delivers the mixed gases to a patient.
- (b) Classification. Class I (general controls). The device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter subject to the limitations in §868.9.

[47 FR 31142, July 16, 1982, as amended at 61 FR 1120, Jan. 16, 1996; 66 FR 38795, July 25, 2001]

§ 868.5610 Membrane lung for longterm pulmonary support.

(a) *Identification*. A membrane lung for long-term pulmonary support is a device used to provide to a patient

extracorporeal blood oxygenation for longer than 24 hours.

- (b) Classification. Class III (premarket approval).
- (c) Date PMA or notice of completion of a PDP is required. No effective date has been established of the requirement for premarket approval. See §868.3.

 $[47\ FR\ 31142,\ July\ 16,\ 1982,\ as\ amended\ at\ 52\ FR\ 17735,\ May\ 11,\ 1987]$

§868.5620 Breathing mouthpiece.

- (a) *Identification*. A breathing mouthpiece is a rigid device that is inserted into a patient's mouth and that connects with diagnostic or therapeutic respiratory devices.
- (b) Classification. Class I (general controls). The device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter subject to §868.9.

[47 FR 31142, July 16, 1982, as amended at 65 FR 2313, Jan. 14, 2000]

§868.5630 Nebulizer.

- (a) *Identification*. A nebulizer is a device intended to spray liquids in aerosol form into gases that are delivered directly to the patient for breathing. Heated, ultrasonic, gas, venturi, and refillable nebulizers are included in this generic type of device.
- (b) Classification. Class II (performance standards).

§ 868.5640 Medicinal nonventilatory nebulizer (atomizer).

- (a) *Identification*. A medicinal nonventilatory nebulizer (atomizer) is a device that is intended to spray liquid medication in aerosol form into the air that a patient will breathe.
- (b) Classification. Class I (general controls). The device is exempt from the premarket notification procedures in subpart E of part 807 of this chapter subject to §868.9.

 $[47\ FR\ 31142,\ July\ 16,\ 1982,\ as\ amended\ at\ 65\ FR\ 2313,\ Jan.\ 14,\ 2000]$

§868.5650 Esophageal obturator.

(a) *Identification*. An esophageal obturator is a device inserted through a patient's mouth to aid ventilation of the patient during emergency resuscitation by occluding (blocking) the esophagus, thereby permitting positive pressure